PCT WOT INTERNATIONAL APPLICATION (51) International Patter Classification 5:

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



WO 93/19007

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCI)

(11) International Publication Number:

C01B 31/02	ΛI	(43) International Publication Date: 30 September 1993 (30.09.93)
(21) International Application Number: PCT/US: (22) International Filing Date: 4 September 1992.(change Place, 53 State Street, Boston, MA 02109 (US).
(30) Priority data: 853,002 - 18 March 1992 (18.03.92) (71) Applicant: MASSACHUSETTS INSTITUTE OI NOLOGY (US/VIS); 77 Massachusetts Aven- bridgs, MA 20135 (US).	F TEC	(81) Designated State: JP. US Published Dith international search report. III-
(72) Insvalous: HOWARD, Jack, B.; 24 Central Str. chester, MA 01890 (US). LAFELUR, Arthur, Wall Stroet, Arlington, MA 02174 (US), QUI Michael, A.; 17 Clearview Drive, Bodford, No. B4A 3C9 (US).	LLIA	16 M.

(54) Tide: METASTABLE FULLERENES

(57) Abstract

Metastable fallerenes have been identified and isolatoi, in particular, the metastable fallerenes are inomers of the stable forms of G₀ and G₀ follerenes. Metastable fullerenes convert to a fallerene of the same molecular weight upon handing. A fulleren incomer is industed and identified. The fullerene incomer is industed and identified for a fullerene having a formula G₀, where s is in the range of 60 to 75 and further having a retention time in liquid chromatography differing from that of the fellerene having the formula G₀.

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Metastable Fullerene s

Cross-Reference to Related Patents

The present application is a continuation-in-part application of copending application U.S.S.N. 07/605,310, filed May 24, 1991.

Field of the Invention

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The present invention relates to closed-caged carbon molecules known as Buckminsterfullerenes or fullerenes and isomers thereof.

Background of the Invention

Pullerenes were first reported by Kroto et al. in carbon vapor produced by laser irradiation of graphite (Wature 318, 162-164 (1985)). Pullerene C₆ is a closed cage carbon structure containing 20 sto-membered rings and 12 fivemembered rings with the appearance of a soccer ball. There has been a surge of scientific interest in these compounds because they represent a new class of carbon in addition to the two known forms, graphite and diamond.

Pollarenes have many potential applications. The ability to intercalate melacities into the structure suggests uses as catalysts in industrial processes. The potassium-fullerene C_{ob} is a superconductor with a T_c of 11 K. The fullerene C_{ob} surface is susceptible to chemical reactions such a hydrogenation and fluorination. Fluorinated fullerenes are expected to be good lubricants. Extensive research is being conducted throughout the United states and the world to develop or discover new and useful fullerenes.

30 Diederich et al. (Science 254, 1768-1770(1991) reported the isolation and characterization of isomeric C_n fullerenes. However, isomeric forms of lower fullerenes are unknown, in particular, C_v, where x is less than 78.

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Summary of the Invention

In the parent application of which this is a continuation-in-part it is disclosed that fullerenes could be produced in flames.

In one aspect of the present invention, the applicants have recognized that, in addition to the fullerene reported in the purent application, sizable quantities of a metastable fullerene can also be present. By "messtable", as that term is used herein, it is meant a species that is transient but of sufficient stability to permit isolation under specific conditions. In preferred embodiments of the hirvestion, the molecular formula of the metastable fullerene is C_w. To other preferred embodiments, the molecular formula of the metastable fullerene coverts to a fullerene having the same molecular formula upon heating. The "isolated peniagon rule" requires that no two peniagonal carbon sub-units of closed-eggs structure lie adjacent to one another. Metastability may be the result of 15 having adjacent peniagonal can onsolven. Metastability may be the result of 15

In another aspect of the invention, a fullerene is more is provided having M* and M* loss identical to that of a stable fullerene and further having a release of the individual control of the fullerene having the formula C_v. By "M* and M* inns', as that term is used herein, it is meant the singly and doubly charged parent ion peaks identified upon mass spectroscopic analysis. By "isomer," as that term is used herein, it is meant a structural or conformational variation of a compound having the same nonlocular formula as the known fullerenes. Somers of C_w. C_w

In a preferred embodiment of the invention, the fullerene isomer is metastable. Isomers of fullerene C₆₀ were considered to be unlikely in view of the fact that the known "soccer ball" structure which possesses 12 five-

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pentagonal rings completely surrounded by the 20 six-membered or hexagonal rings is the only possible arrangement which obeys the "isolated pentagon rule".

In another aspect of the invention, the yield of a metastable isomer can 5 be optimized in any fullerene source by rapidly quenching the fullerenes at a location in the process where the fullerenes are being produced. In a preferred embodiment, metastable isomer yield is improved by avoiding high temperatures in post-production treatments. The metastable fullerene converts readily under the high temperature conditions of the flame. By 10 quenching the reaction before there is sufficient time to allow the complete conversion of the metastable to the stable fullerene, the yield of metastable fullerene is enhanced. Likewise, means taken to avoid subjecting the metastable compound to conditions that would encourage conversion to the more stable form of the fullerene during isolation and purification will 15 enhance the final yield of metastable fullerene. Means of quenching include, but are not limited to, inserting a collection tube near the location of fullerene formation into which is introduced the cold nitrogen evaporant from liquid nitrogen source or by injecting a fluid from high velocity jets into the location of fullerene production.

Analysis of materials made by the process disclosed in the parent application indicate the presence of isomers of C_{so} and C_{ro} fullerenes. In the purification of soot samples by high performance liquid chromatography (HPLC), species were observed with retention times differing from those of the Cs and Cn fullerenes. Mass spectrometer analysis of the different species 25 indicated that they had the same mass as the known C₆₀ and C₇₀ fullerenes. It is speculated that the isomer of the Co fullerene may have a flattened, pillow-like configuration, a cylindrical structure or even a flattened cylindrical structure. It has also been determined that the isomers are metastable and will revert to the respective fullerene under appropriate 30 conditions. Because of their metastable nature and because of their different structure, these isomers may be more useful than the base fullerene for certain application. For example, the metastable isomer may be more easily WO 93/19007

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doped with atoms such as potassium for making a known superconducting fullerene. Another potential application of the metastable isomer is as a starting material for making diamonds using the application of extremely high pressures, for example, diamond anviis. These meta stable fullerenes 5 may be useful as starting materials for chemical addition or polymerization reactions.

Brief Description of the Drawing

In the Drawing:

10 Figure 1 is a high performance liquid chromatogram (HPLC) of a toluene extract of a flame soot indicating C₁₀ and C₂₀ as well as new peaks A-D;

Figure 2 is selected ion chromatograms (SIC) from the HPLC-MS analysis of a flame soot;

Figure 3 is a background-subtracted positive mass spectra obtained at the crests of chromatographic peaks annotated in Figure 2;

Figure 4 is a background-subtracted negative ion mass spectra obtained at the crests of chromatographic peaks annotated in Figure 2; and Figure 5 illustrates the time-dependence of chromatographic peak

20 areas from HPLC analysis of flame-derived fullerene extract in boiling toluene under argon.

Description of the Preferred Embodiment

The structure of the remarkably shible carbon cluster Cap first discovered by Kroto et al. (Nature 318, 162 (1988)) in vapor from laser tradiation of graphite and subsequently produced in macroscopic quantities by resistive heating of graphite under an inert atmosphere (Kristchmer et al., Nature 387, 384 (1990)). Taylor et al., I/ES Chem. Cammun. 1423 (1990), Aja et al., I, Flys. Chem. 94, 8630 (1990). Hautler et al., I, Flys. Chem. 94, 8634-8636 (1990), is now widely accepted as a truncated icosahedron ("soccer ball fullerme"), while the homologius C_p in assigned an ellipsoidal cage structure Crusby ball; Stoddart, August. Chem. Int. Ed. Engl. 30, 70-72 (1991).

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"Buckminsterfullerene" and "fullerene" as these terms are used herein refer to these highly symmetrical closed-cage carbon clusters. The parent application discloses that Cas and Cas fullerenes, isolated from samples of condensible compounds and soot collected from controlled combustion of benzene in pre-mixed laminar flames, were spectroscopically indistinguishable from fullerenes prepared from graphite. Analysis of the flame samples by high-performance liquid chromatography (HPLC) with ultraviolet spectroscopic detection (UVD) also revealed the presence of several additional components with fullerene-like characteristics.

Efficient methods for the analysis of polycyclic aromatic compounds (PACs) of high molecular weight, by HPLC or supercritical fluid chromatography (SFC) coupled directly to mass spectrometry (LC-MS or SFC-MS), have been developed recently. These methods employ either a movingbelt interface with flash heating of the analyte as it enters the EI source, or 15 a heated pneumatic nebulizer coupled to an atmospheric pressure chemical ionization (APCI) source in which electron transfer to benzene molecular cations is arranged to be the dominant ionization mechanism. In the present work the flame soot samples were analyzed by these complementary LC-MS techniques, as well as by HPLC interfaced on-line to a diode array detector 20 for UV spectroscopy.

Experimental.

Materials. The extracts of flame-generated soots and condensates were identical to those described in the parent application. A sample of soot produced by resistive heating of graphite, and one of a purified extract of Co 25 from such soot (containing also a few per cent of the C_m fullerene), were obtained from the Texas Fullerenes Corpn., Houston, Texas 77030.

HPLC with UV spectroscopy. The column used was 25cm long x 2.1 mm i.d., with Vydac 201TP C18 packing. The initial solvent was 100% acetonitrile for 5 min, then programmed linearly to 100% dichloromethane 30 over 45 min, held for 5 min, then programmed back to initial composition over 5 min. The mobile phase flow rate was 200 µL/min, with an injection volume of 5 ul., A HP1090M liquid chromatograph (Hewlett Packard 5

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Co.,Palo Alto, CA, USA), equipped with a binary DR5 solvent delivery system, a HP1040A diode array detector, and a HP7994A data system, was used in all LC-UV analyses. The detector was configured for continuous full UV scan acquisition (220-600 nm).

HPLC with on-line mass spectrometry. The HPLC conditions were identical to those used in the LC-UV analyses. In one set of experiments the HPLC efficient (no split) was introduced to an electron ionization mass spectrometer (VG 20-250 quadrupole mass filter, VG Mass Lab, Altrincham, U.K.) via a VG moving belt interface. The conditions used were similar to 10 those for LC-MS analysis of PACs with molecular weights of up to 600, except that the fullerenes and related compounds required higher temperatures for efficient volatilization. The nominal electron energy was 70 eV, with a trap current of 100 μA. The source temperature was 350°C, and the flash belt heater at the tip of the belt interface, located within the body 15 of the ion source, was operated at maximum power (belt surface temperature unknown). In addition, the belt clean-up heater (also at maximum power) and wash-bath had to be employed in order to avoid memory effects. A VG 11-250I data system was used for instrument control and for data acquisition and processing.

The LC-MS experiments employing electron transfer APCI were conducted using an API III triple quadrupole instrument (SCIEX, Thornhill, Ontario), equipped with a heated pneumatic nebuliser interface (SCIEX). The pneumatic nebuliser is contained within a concentric quartz heating tube, itself located within the room temperature APCI source; the indicating 25 thermocouple is located on the exterior surface of this heating tube, together with the heating element. Experiments conducted on this nebuliser interface mounted on the bench but under conditions otherwise identical to those used in the LC-MS analyses, with a second thermocouple used to probe the gas temperatures within the interface, showed that the maximum temperatures 30 experienced by the analytes under these conditions fell in the range 80-100°C. Medical quality air was normally used as the nebulizing gas, but was replaced by high-purity nitrogen for a few experiments; high purity nitrogen was always used for the counter-current gas flow (the "gas curtain") which prevents unevaporated droplets and other debris from clogging the orifice connecting the API chamber to the mass spectrometer vacuum system.

Ionization was achieved in a fashion very similar to that described for SFC-MS analysis of PACs. The APCT plasma was sustained by a cold corona discharge featurels set est need manistant et at SIV. Introduction of benzene vapor vis the nebulizing gas lake ensured that the dominant positive reactant ions in the plasma were CH₂*, together with some water cluster lons from residual water in the system. Under these conditions the predominant io ionization mechanism for polycyclic aromatic hydrocarbons (PAEs) involves electron transfer to CH₂*, with some tendency for protonation. Characterization of this APCI plasma in the absence of analytes also showed a significant population of benzene-derived ions at m/z 91, presumably CH₂* long to row/limits structure.

Results and Discussion.

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Admitifaction of Pails 1 and 11 as closed-coge fullerents. The LC-UV analysis of the flame-scot extract (Fig. 1) was compared to that of an extract of soot produced by resistive heating of graphite. This comparison permitted 20 identification of peaks 1 and 11 in Fig. 1 as C.g. and C.g. fullerents, wis both retention times and UV spectra. Further confirmation was obtained by HPIC-MS experiments using a moving-belt interface and an El source. The reconstructed total ion chromatogram (not shown) matched Fig. 1 extremely well, although some of the smaller HPIC peaks were not well defined. The El El mass spectra (not shown) recorded at the crests of peaks 1 and 11 (Fig. 1) were identical to those for C.g. and COd.

In the striffication of Pauls B and C as C_m and C_m species. Peaks B and C in Fig. 1 were of particular interest since then were absent from the chromatogram of extract of spoot from resistive heating of graphite. As shown in Fig. 1 the UV spectra obtained at the crosts of Peaks B and C differ from those of the authentic C_m and C_m fullerenses (Peaks I and III), but obtained with those of the authentic C_m and C_m fullerenses (See pasks I and III), but obtained with the cross the spectra acquired in full full remership (See particularistics). Surprisingly, the El mass spectra acquired in

the moving-helt LC-MS experiments at the cross of peaks B and C were identical to those for peaks I and II respectively, showing intense M^* and M^* ions with appropriate isotopic distributions. This combined, IVV and EI evidence strongly suggested that peaks B and C arose from isomers of the C_m 5 and C_m fullerenes.

However, at least two other interpretations of these HPICAMS experiments are possible, based on the hypothesis that the additional abundant components in the flames not extract (eases B and C) were adducts (e.g. with oxygm) of the C_a and C_p, fullerenes. Such compounds could 10 conceivably yield negligible intensities of molecular ions upon electron ionization, but abundant fragment ions C_a* and C_p*. However, the intense dications C_a* and C_a* are more difficult to rationalize on this basis. Alternatively, thermal decomposition of labels fullerene adducts to the parent compounds, subsequent to dromatographic separation and during the toward the control of the composition of the control of the

This hypothests of labile fullerens adducts, which survived the chromatography but decomposed either before or after II, was tested view treatment of each of the soot extracts (sgenerated in flames or by resistive to the stream of cach of the soot extracts (sgenerated in flames or by resistive beating of graphite) with ferrous ammonium sulphaste, and separately with sodium borohydrife, in order to reduce suspected coygen adducts. In other experiments the extracts were treated by passage through an activated alumina column in order to remove any hydroperoxides. All three treatments left the chromatogram (flg. 1) unchanged. However, refluxing the extracts in boiling tolunes (III'C) under argon resulted in a slow loss (balf-life approximately 1 hour) of peaks B and C, with corresponding increases in peaks I and II whose UV spectra were thereby unchanged. A systematic study of the time dependence of these thermal transformations was undertaken, but this is best described below, after discussion of the 30 confirmation of the leftenties of the HPLC peaks.

Clearly, a gentler and more sensitive method of HPLC-MS coupling was required. This was achieved using an APCT source, similar to that used

for SPC-MS analyses of PACs. The HPLC effluent was nebulized to micronsized droplets within a heated quartz tube; the maximum gas temperature in this region was about 110°C in the first experiments, later reduced to 80-90°C with no significant change in the results obtained. The mobile phase 5 was rapidly evaporated from these droplets via interactions with the heated gas (nitrogen saturated with benzene vapor, together with vaporized mobile phase), leaving extremely small particles each containing only a few analyte molecules. The vapor pressure of such small particles can be many times larger than that of the bulk material, permitting vaporization of analytes 10 which are thermally labile when heated in the bulk phase. The vaporized sample was then drifted into a cold corona-discharge plasma, whose composition was controlled by manipulating that of the atmosphere in the APCI source. In this work post-column addition of benzene ensured that the dominant mechanism for formation of positive ions from the sample was 15 electron transfer to C_eH_e* ions, whose recombination energy is 9.25 eV. Since the ionization energy of $C_{eq}(g)$ is about 7.8 eV, electron transfer ionization was efficient and the excess energy of 1.5 eV was readily degraded by collisions with the atmosphere. A minor ionization mechanism was formation of adducts with tropylium ions (C,H,*, m/z 91) formed from 20 benzene in the APCI plasma. The sample-derived ions thus transmitted to the mass analyzer were internally cold, yielding mass spectra which exhibit only molecular and adduct ions with little or no fragmentation.

Results obtained by such HFLC-MS analysis of the flame sample are summarized in Fig. 2 and 3. The selected ion chromatogram (SIC) for C₂ at n/z 720 (Fig. 2a) shows two peaks at retention times matching those of peaks B and I (Fig. 1). The APCI mass spectra recorded at these two peak crests were identical to one another (Fig. 3a), dominated by molecular ions (a.* although the isotopic intensity distributions indicate a minor contribution from C₂H* formed by proton transfer from residual water cluster ions. Corresponding data acquired for the extract of the graphite soot (not shown) gave an identical spectrum for peak I (C₂ fullerene). Analogous results obtained for peaks C and II (Fig. 2 c and 3b) suggest that peak C is

a C_{70} isomer and that other C_{70} isomers are present at lower abundances.

The short contact times (< 5 sec) and low temperatures (109°C and 8090°C) experienced by the analytes in the APC interface, compared with those
in the refluxing folione experiments, make it improbable that peaks A and
5 B correspond to thermally labile adducts of the fullereness which survived the
chromatography but decomposed to C₆ and C₉ in the heated nebuliser. This
conclusions was verified by flow-injection of an aliquot of the flame soot
extract through the heated nebuliser, using the same mobile phase and
thermal settings as for the HPIC-NAS experiments but physically removed
of from the mass performents and corona discharge. The emerging wapor was
condensed on a cold surface, redissolved, and analyzed. All of peaks A-D
survived this treatment, confirming the stability of the sample under these
HPIC-NAS conditions price to lonization.

It is difficult to produce equally convincing evidence to contradict the 15 hypothesis of fullerene adducts which survive both the chromatography and vaporization in the heated nebuliser interface, but whose molecular ions undergo facile and complete fragmentation to C_{es}^{-} and C_{gs}^{+} . However, in our hands this ionization method invariably produces abundant molecular ions, with no fragmentation, from fragile and relatively involatile analytes of 20 a wide variety of structural types. In addition the evidence for formation of (Ca+H)* and (Ca+C4H2)* ions, and analogous adduct ions from C20 (Fig.s 3a and 3b), support the contention that peaks B and C correspond to C, and C, species, respectively. Further support of this interpretation is provided by negative ion APCI spectra (Fig. 4), obtained under LC-MS conditions identical to those used for the positive ion work (Fig.s 2 and 3) but omitting the benzene. Under these conditions, ionization of the Cs and Cn compounds proceeds via two principal mechanisms, viz. electron attachment and chloride attachment (the Cl' arising from the dichloromethane in the HPLC mobile phase): As shown in Fig. 4 the spectrum obtained for Peak B is almost 30 identical to that for Peak I (the authentic Cas fullerene), further supporting the present contention that Peak B corresponds to a chromatographically distinguishable form of C₆₀ rather than to some adduct of C₆₀ which, WO 93/19007 PCT/US92/07491

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subsequent to ionization, fragments immediately to yield a mass spectrum which is identical to that formed directly from ionization of the authentic fullerene. Slight differences between the two spectra in Fig. 4 are due mostly to differences between the compositions of the APCI plasma in the two cases, 5 under the HFIC gradient conditions used in these LCMS experiments. Detailed interpretation of these negative ion spectra will be published separately, but it is noted here that contributions to the spectra include those from the nebulizing gas dissually air) and from redistal water. Finally, a 10 comparison of the analogous negative ion spectra (not shown), obtained for Peaks C and II, similarly supported the proposal that both peaks correspond to Cs. spectra

The evidence supports the contention that Peaks B and C, in the HPLC
chromatograms of the flame-generated fullerenes, arise from isomers of the
15 C₀₀ and C₂₀ fullerenes, respectively.

Identification of other HPLC peaks as higher carbon clusters or monoxides. Higher carbon clusters were also observed at lower abundances in the flammagnerated extracts. Thus, peak D (Fig. 1) corresponds to C_{∞} (Fig. 2a). The two minor non-amostated peaks preceding Peak D (Fig. 1) correspond to C_{∞} (Fig. 2a and 3c), a chiral form of C_{∞} was isolated and characterized very recently by Diedericht d d. Evidence was also obtained (not shown) for several forms of C_{∞} and C_{∞} present at very low abundances in this extract. Note that the mass spectra obtained for these minor components, e.g. Fig. 3c, were obtained near the ion statistical limit so that significant random 25 deviations of the isotope intensity distributions, from those predicted from assumed monocular formulae. were observed.

Peak A (Fig. 1) was identified as a monoxide C_nO six the AFCI mass spectrum, which showed evidence for increased importance of protonation to form (04 + H)* (Fig. 3d) relative to that of the C_n compounds. The SiC for 30 m/z 736 (Fig. 2b) shows evidence for at least 5 chromatographically distinguishable species giving rise to C_nO*, but two of these have retention times identical to those of peaks B and I and were probably formed as

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oxidation artifacts of the corresponding carbon clusters in the APCI plasma. This was confirmed by repeating the analysis with meticulous exclusion of air (using high purity nitrogen as nebulizing gas) and of water, insofar as this was possible; as a result the intensities of the oxidation artifacts were reduced 5 by a factor of about 5, while those of the chromatographically distinguishable monoxides remained unaffected. Analysis of the graphite soot gave an SIC for m/z 736 (not shown) containing only peak A plus a weak APCI artifact corresponding to Peak I. These observations are susceptible to an interpretation in which Peak A is the unique monoxide of Cao 10 buckminsterfullerene (all carbons equivalent), while the other non-artifact signals in Fig. 2b correspond to monoxides of less symmetrical Con isomers not present in the graphite soot. Fig. 2d shows similar evidence for several CnO isomers in the flame soot, apart from the APCI artifacts. The corresponding experiment on the graphite soot extract (not shown) showed 15 evidence for only two C_nO compounds apart from the APCI artifact at a retention time corresponding to Peak II; this observation is again qualitatively consistent with the requirement that the new C20 isomers (Fig. 2c) possess a lower degree of molecular symmetry than does the authentic fullerene. One C.O compound has been isolated previously14 from a soot produced by resistive heating of graphite.

All of these findings summarized in Fig.s 2 and 3 were qualitatively confirmed by repeating the HPLC-MS analyses in negative ion mode. The observed multiplicities of the C60O and C70O isomers provide circumstantial evidence supporting the conclusion, described above, that Peaks B and C 25 correspond to isomers of the C₁₀ and C₂₀ fullerenes with lower degrees of molecular symmetry.

Rates of thermal conversion of C_{so} and C_{70} isomers to stable fullerenes. The lower stabilities of the newly observed isomers, relative to those of the respective previously observed fullerene structures, is qualitatively consistent 30 with calculated relative stabilities by Schmalz et al. (J. Am. Chem. Soc. 110, 1113 (1988)) and with more recent predictions by Goeres and Sedimayr (Chem. Phys. Lett. 184, 310 (1991)) based upon a theoretical nucleation WO 93/19007 PCT/US92/07491

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mechanism for formation of fullerenes.

In the present work, the rate of thermal conversion of the new isomers to their fullerene counterparts was studied in solution in deserated botting tolones (11°C), in an argon atmosphere. A few IC-MS experiments, using 5 the positive ion AFCI method, were run to confirm the identities of reactinals and products of the thermolysis process, but the quantitative data were obtained using IC-UV analyses similar to that illustrated in Fig. 1. In all cases coroneme was added to the extract clustions as a non-reactive internal standard, to avoid concentration errors due to evaporation of solvent.

10 Several experiments were run in parallel, whit quenching at varying reaction times. The results thus obtained are illustrated in Rg. 5, for the C_x species only (similar results were obtained for the C_x species). The peak areas plotted in Fig. 5 are for appropriate HFIC peaks annotated in Fig. 1, monitored with the integrated optical absorptions over the range 269-390 nm; 15 thus, no correction was possible for differences in molar extinction coefficients.

The decay of Peak B (new C_n isomer) is observed to occur with a halflife of approximately 1 hour. The corresponding rise of Peak I (authentic C_n fullerene) does not compensate for the loss of Peak B intensity, as is most 20 readily seen from the plot (Big. 5) of the sum of the two intensities as a function of time. This could be the in part to a higher integrated molar absorption for the new isomer (Peak B) than for its fullerene counterpart, or to possible side-reactions. A small but significant rise (Big. 5) in the intensity of Peak A (C_mO) suggests that some degree of oxidation occurred despite the 25 precautions taken. Also, other processes such as polymerization would have yielded side-products which would not have been detected in these experiments.

What is claimed is:

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. A metastable fullerene.

- The metastable fullerene of claim 1 wherein the structure of the fullerene includes at least one pair of adjacent pentagonal carbon units.
- 3. The metastable fullerene of claim 1 having the molecular formula of C_{nor}
- The metastable fullerene of claim 1 having the molecular 10 formula of C₇₀.
 - The metastable fullerene of claim 1 wherein the metastable fullerene is converted to a fullerene of same molecular formula upon heating.

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- A fullerene isomer, characterized by
- a M^{\star} and M^{2r} ions by mass spectroscopy identical to that of a stable fullerene; and
- a retention time in liquid chromatography differing from the 20 $\,$ retention time of the fullerene having a formula C_{rr}
 - A fullerene isomer, characterized by
 - a M^{+} and M^{2+} ions by mass spectroscopy identical to that of a fullerene:
- 25 a retention time in liquid chromatography differing from the retention time of the fullerene; and
 - which has a structure includes at least one pair of adjacent pentagonal carbon units.
- 30 8. The fullerene isomer of claim 6, wherein the isomer is metastable.

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- The fullerene isomer of claim 6, the isomer being a structural isomer of the fullerene having a structure of C_x.
- 10. The fullerene isomer of claim 6, the isomer being a 5 conformational isomer of the fullerene having a formula of C_c.
 - 11. The fullerene isomer of claim 6 or 7 wherein the molecular formula is $C_{\mbox{\tiny MF}}$
- 10 12. The fullerene isomer of claim 6 or 7 wherein the molecular formula is C₇₀.
 - 13. The fullerene isomer of claim 6 or 7 wherein the molecular formula is C_{α}

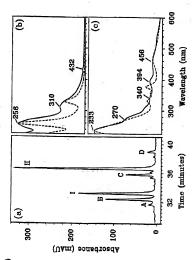
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- 14. The fullerene isomer of claim 6 or 7 wherein the molecular formula is C_{64}
- $15. \quad \text{The fullerene isomer of claim 6 or 7 wherein the molecular} \\ 20. \quad \text{formula is C_{60}}.$
 - 16. The fullerene isomer of claim 6 or 7 wherein the molecula formula is C_{st} .
 - The fullerene isomer of claim 6, wherein the structure of the isomeric fullerene includes at least one pair of adjacent pentagonal carbon units.
- 18. The method of optimizing the yield of a metastable fullerene 30 isomer in a fullerene source, comprising increasing the quench rate of fullerenes at a point where fullerenes are being made.

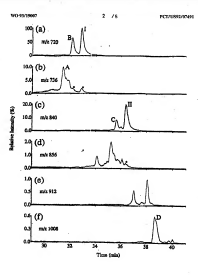
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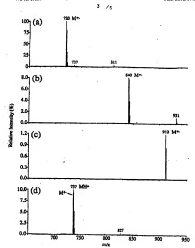
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 The method of claim 18, further including avoiding high temperatures in a post-production treatment of a collected product.



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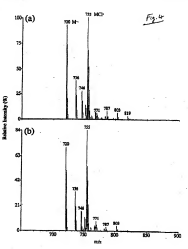
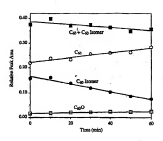


Fig. 5



INTERNATIONAL SEARCH REPORT

			Internacional Application No	PCT/US 92/07491		
I. CLASSII	ICATION OF SUBJ	CT MATTER (if several classification	symbols apply, indicate all)*			
Int.C		C 01 B 31/02	Classification and IPC			
W 0001 P.C	SEARCHED					
II. FIELDS	SEARCHED	Marie Per	estation Searched			
Classificat		Potition Document	Classification Symbols			
Cammon	134 System		Classicator symbols			
Int.C1	1.5	C 01 B 31/00				
		Documentation Searched other to the Extent that such Documents	r than Minimum Documentation are Included in the Fields Sourched *			
III. DOCU	MENTS CONSIDERS	D TO BE RELEVANT?				
Category *		ocument, 12 with indication, where approp	rists, of the relevant passages 12	Relevant to Claim No.13		
Α	Chemical Moximusts, vol. 115, no. 14, 7 October 1991, (Columbus, Ohio, US), 1, COULDMENE et al.: "Study of 74 CGU carbon isomer aggregates by the Husckeln enthod", see page 455, abstract no. 1428389, 8, J. CHIM. PMYS. PMYS.—CHIM. BIUL. 1991, 88(5), 867-74					
٨	1992, "Struc isomer	al Abstracts, vol. 116 (Columbus, Ohio, US), ture and stability of s", see page 546, abst REV. B: CONDENS. MATTE 6	L. GOODWIN: some fullerene C60 ract no. 67712r, &			
	a categories of cited do		"I" later decement pelitabed after	the international filing date		
A 60	cament defining the go	neral state of the art which is not alar reiovance	"I" later document published after or priority date and not in con- cised to understand the principl	in or theory underlying the		
		lished on or after the incernational				
77.44	ing date	m device on miletin chiledia or	"A" document of particular relevan- caused be considered novel or o involve an inventive step	mazet be considered to		
whi	ich is cited to establish	or équites on priority claim(s) or the publication date of another eason (at specified)	"> document of particular relevan- cassot be considered to involve document is combined with one ments, such combination being	or the claimed invention		
		erui disciosure, use, exhibition or	equant be considered to involve document in combined with one	an inventive step when the or more other such docu-		
043	PER MICHEL		ments, such combination being	coveres to a person skilled		
lac.	or than the priority da	to the international filling date but to claimed	"A" document member of the same	patent foresty		
IV. CERT	FICATION					
		the Leserational Search	Date of Mailing of this Internal	ional Search Report		
	29-12-	1992	18. 01. 93			
Internations	ai Searching Authority		Signature of Authorized Offices			
		AN PATENT OFFICE	J. BREBION			
New PCTASA	J230 (second short) Limon	ry 19830				

International Application No. Page 2 PCT/US 92/07491

		/US 92/07491
	TIS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET) Clusters of Document, with Indication, where appropriate, of the relevant possesses	Relevant to Claire No
Category *	Citation of Doctatesti, with their catego, waters appropriate, or the postwar products	Accessed to Course No.
A	Nature, vol. 352, 11 July 1991, (London, GB), J.B. HOWARd et al.: "Fullerenes C60 and C70 in flames", pages 189-161, see page 140, left-hand column, paragraph 1; flyure 2	,
т	Chemical Physics Letters, vol. 184, no. 4, 27 September 1991, (Masterdam, NL), A. GOEMES et al.: "On the nucleation mechanism of effective fullerite condensation", pages 310-317 (cited in the application)	
T	Journal of the American Chemical Society, vol. 110, no. 4, 1988, (Washington, DC, US), T.G. SCHMALZ et al.: "Elemental carbon cages", pages 1113-1127 (cited in the application)	
P,X	Chemical Abstracts, vol. 117, no. 8, 24 August 1992, (Columbus, Ohito, US), J.B. MCMARD et al.: "Production of CSG and C70 fullerness in benzeme-oxygen flames", see page 356, abstract no. 82176h, & J. PHYS. CHEM. 1992, 95(16), 6657-62	1,3,4
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